Noninvasive Index of Cryorecovery and Growth Potential for Human Follicles In Vitro

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Summary: Cryopreservation of individual follicles ensures the maintenance of TZPs, which are key to the survival and growth of secondary follicles post thaw.

Abbreviations: OTC-ovarian tissue cryopreservation; TZP-transzonal projections; IFM-in vitro follicle maturation

Key Words: transzonal projections, follicle, cryopreservation, birefringence, alginate, fertility preservation

Abstract
Cryopreservation of oocytes and embryos is commonly used to preserve fertility. However, women undergoing cancer treatment may not have the time or may not be good candidates for these options. Ovarian cortical tissue cryopreservation and subsequent tissue transplant has been proven successful yet inefficient in preserving larger secondary follicles, and is not recommended as a fertility preservation option for women with certain cancers. We evaluated cryopreservation of individual follicles as an alternative option in rodents, non-human and human primates. Under optimal conditions, cryopreserved mouse secondary follicles were able to re-establish granulosa cell-oocyte interactions, which are essential for subsequent follicle growth. Individual secondary follicles survived cryopreservation, were able to be cultured in a 3D alginate hydrogel matrix to the antral stage, and the enclosed oocytes were competent for fertilization. Using a vital imaging technique (pol-scope) employed in many fertility centers, we were able to bioassay the thawed, cultured follicles for the presence of transzonal connections between the somatic and germ cells. Perturbations in these linkages were shown to be reversed when follicles were cryopreserved under optimal freezing conditions. We applied the optimized cryopreservation protocol to isolated rhesus monkey and human secondary follicles, and using the birefringent bioassay, we were able to show good correlation between early follicle growth and healthy somatic cell-oocyte connections. Our results suggest that ovarian follicles can be cryopreserved, thawed and analyzed non-invasively making follicle preservation an additional option for young cancer patients.
Introduction

Approximately 39 in 100,000 women under the age of 44 were diagnosed with cancer between 2002-2006, and of these women, 84% had a favorable prognosis (NCI SEER cancer statistics 2006). Unfortunately, many cancer therapies, such as alkylating agents and pelvic radiation, render women sub-fertile or infertile by damaging growing follicles and depleting follicular reserves [1]. Prior to cancer therapy, current fertility-sparing options for women include oocyte and embryo cryopreservation; however, women must delay treatment to undergo hormonal stimulation to obtain enough oocytes for in vitro fertilization (IVF) or cryopreservation, and hyperstimulation may not be possible for women facing hormone-sensitive cancers or for young girls [2, 3]. Ovarian tissue cryopreservation (OTC) is offered as an experimental option for fertility preservation prior to cancer treatment [4]. As of 2009, six babies have been born as a result of autotransplantation of cryopreserved tissue [5-8], and several more have been born after non-cryopreserved or frozen heterologous transplants between monozygotic twins [9-11].

Successful OTC transplants use the outer 1 mm of ovarian cortex, which is rich in the immature primordial follicles that comprise the pool of follicles that enter the growing population over time. Unfortunately, the larger follicles, which include primary, secondary and multi-layered secondary follicles, are either discarded prior to freezing or are destroyed in the cryopreservation/vitrification process [12-14]. Moreover, ovarian tissue collected before treatment may contain residual cancer cells, thus, cryopreservation of these immature follicles could be used either in transplant to minimize disease or used for in vitro follicle growth with the goal of providing a mature gamete for fertilization.

Several studies have shown that either cryopreservation or vitrification of individual mouse follicles has resulted in live births; however, these cryopreservation protocols have not been tested on primate or human tissue [15-17]. Moreover, culture systems used to grow mouse follicles post thaw are two-dimensional systems [18] or three-dimensional culture systems [19, 20] that fail to support human follicle growth or antrum formation over an extended period of time. Previous reports in the mouse [21], nonhuman primate [22] and human [23] have shown that in-follicle maturation (IFM) in a 3D alginate-hydrogel matrix is capable of supporting follicle growth to generate mature oocytes competent for fertilization in vitro [21, 23]. The alginate hydrogel culture system provides an excellent tool for determining whether follicle and oocyte growth can be supported following individual follicle cryopreservation and thawing.

One of the challenges faced by the field is the development of tools that can assess follicle health immediately after cryothaw. Moreover, since human tissue is so rarely available for research, robust, non-invasive, vital technologies are urgently needed. Presently, in many fertility clinics the use of the pol-scope is used to determine spindle location prior to intra-cytoplasmic sperm injection [24, 25] particularly after oocyte cryopreservation [26]. This technique also aids in spindle enucleation in the process of cloning [27]. Interestingly, the pol-scope has also been used to assess the thickness of the zona pellucida in hamster embryos during assisted hatching [28]. We too elected to use the anatomically rich region of the zona pellucida as an opportunity to solve each of these methodological problems. The germ cell and the surrounding somatic cells must communicate throughout development and do so through transzonal projections (TZPs), that extend through the glycoprotein rich zona pellucida and connect via gap junctions on the oocyte membrane. TZPs are essential for the bidirectional transport of molecules such as amino acids [29, 30], cAMP [31, 32], glucose [33] and pyruvate [34], which are essential for oocyte
growth. Starting in the primordial follicle stage, TZPs form from initial, simple cell-cell gap junctions that are continuously remodeled and elongated during the formation of the zona pellucida and the course of oocyte growth [35]. Oocyte and follicle growth are dependent on the maintenance of these structures for survival, but due to their composition of F-actin and microtubules [36], they are very sensitive to changes in temperature and shear stress, and are vulnerable during cryopreservation [12]. The TZP network is predicted to be a good biomarker of follicle health, since it is necessary for bidirectional communication between the somatic cells and the oocyte. Here, we assayed the quality of these connections in non-cryopreserved and cryopreserved rodent, non-human primate, and human secondary follicles over time. We assessed the utility of a non-invasive microscopic technique (pol-scope), currently used in several clinics, for providing an in-depth measure of TZP integrity. Since destructive technologies are not suitable for application in the human setting, this bioassay provides a powerful new way to assign an early, non-invasive index of cryorecovery and growth potential for human oocytes post thaw.

Materials and Methods

Animals
Immature follicles were isolated from prepubertal, 12- to 14-day-old female F1 hybrid (C57BL/6J x CBA/Ca) mice for both non-cryopreserved and cryopreserved experiments. Sperm was prepared from proven CD1 male breeders. Animals were purchased (Harlan, Indianapolis, IN), housed in a temperature- and light-controlled environment (12 L:12 D) and provided with food and water ad libitum. Animals were fed Harlan Teklad Global irradiated 2919 chow, which does not contain soybean or alfalfa meal and therefore contains minimal phytoestrogens. Animals were treated in accordance with the NIH Guide for the Care and Use of Laboratory Animals, and protocols were approved by the IACUC at Northwestern University.

Isolation and growth of mouse follicles
Early secondary follicles (two to three granulosa layers) were isolated from 12- to 14-day-old mice and encapsulated into sterile 0.5% (w/v) alginate beads as described previously [37]. Ovaries were isolated into pre-warmed collection medium (Liebovitz L-15; Invitrogen Corporation, Carlsbad, CA) containing 1 mg/ml bovine serum albumin (BSA) and 50 IU/ml penicillin/streptomycin (Invitrogen). For the cryopreserved follicle group, follicles were mechanically isolated using insulin gauge needles and were stored in L-15 medium prior to cryopreservation as described below. Upon thawing, follicles were stored in maintenance medium, a-MEM containing 1 mg/ml BSA and 50 IU/ml penicillin/streptomycin at 37°C, 5% CO₂ for 2 hours prior to encapsulation in alginate hydrogel according to previously published methods [21]. Follicles were grown in a-MEM containing 1mg/ml bovine fetuin (Sigma-Aldrich, St. Louis, MO, USA), 5µg/ml insulin, 5µg/ml transferrin, 5ng/ml sodium selenite (Sigma-Aldrich, St. Louis, MO, USA), 3mg/ml BSA and 10mIU rhFSH (Gift from Organon, Roseland, NJ, USA).

Isolation and growth of non-human primate follicles
Rhesus macaque ovaries (age 13 years) were obtained from the laboratory of Dr. Leo Towle in the Department of Neurology at the University of Chicago under an approved material transfer agreement. All follicles in this study were collected from non-cryopreserved tissue, which was cut into approximately 1-mm³ pieces in dissection medium (L-15 medium [Invitrogen])
supplemented with 1% serum protein substitute [SPS, CooperSurgical, Inc., Trumbull, CT], 100 IU/ml penicillin, and 100 mg/ml streptomycin [Invitrogen]). The tissue was then enzymatically digested by transferring it into dissection medium containing 0.2% collagenase and 0.02% DNase (Worthington Biochemical, Lakewood, NJ) for 30 minutes at 37°C in a shaking water bath. Follicles were mechanically isolated from the tissue strips using 25-gauge needles, transferred to maintenance medium (aMEM [Invitrogen] supplemented with 1% SPS, 100 IU/ml penicillin and 100 mg/ml streptomycin), and placed in an incubator at 37°C and 5% CO₂ for 2-6 hours prior to encapsulation (non-cryopreserved controls) or cryopreservation. Follicles were grown in a-MEM containing 1mg/ml bovine fetuin (Sigma-Aldrich, St. Louis, MO, USA), 5µg/ml insulin, 5µg/ml transferrin, 5ng/ml sodium selenite (Sigma-Aldrich, St. Louis, MO, USA) and 3mg/ml BSA. No FSH was added to these cultures.

Isolation and growth of human follicles
Portions of four patients’ ovarian tissue (never more than 20% of the total) were donated for research following informed consent under an IRB-approved protocol at Northwestern University. All follicles included in this study were collected from non-cryopreserved tissue, which was cut into approximately 1-mm³ pieces in dissection medium (L-15 medium; Invitrogen) supplemented with 1% SPS (CooperSurgical), 100 IU/ml penicillin, and 100 mg/ml streptomycin (Invitrogen). Human follicle isolation and culture is followed exactly as rhesus follicle isolation an culture at this point.

Cryopreservation and thawing of follicles
After individual follicles were isolated from ovarian tissue, they were washed in collection medium 2 times before transferring to 1 ml cryopreservation medium (~30 follicles per tube) containing L-15 medium, sucrose 35 mg/ml (final concentration), Quinns Advantage serum protein substitute (SPS, CooperSurgical) 10% (v/v) and ethylene glycol 8.83% (v/v) for 30 minutes at room temperature followed by 30 minutes at 4ºC. Follicles were then transferred to cryotubes containing 1 ml of pre-chilled cryoprotectant for 10 minutes.

The tubes were then transferred to a 4ºC freeze control Cryobath (Cryologic, Victoria, Australia). Cryopreservation was controlled by CryoGenesis V software (CryoLogic), which cooled the tubes to -7ºC at a rate of 2ºC/minute. Cryoseeding was performed at -7ºC (permissive temperature; the ideal temperature for crystallization seeding for secondary follicles in ethylene glycol) or -9ºC (non-permissive/negative control), followed by chilling to -40ºC at 0.3ºC/minute and then storage in N₂ (l) until thawing.

For thawing, tubes of follicles were removed from N₂ (l) and held at room temperature for 30 seconds prior to being submerged in a 37ºC waterbath for 2.5 minutes. Contents of the cryotube were removed using a trimmed 1 ml pipet tip and placed in a 60-mm culture dish. Retrieved follicles were transferred to room temperature L-15 medium containing 10% SPS and sucrose 35 mg/ml for 3 x 10 minutes with decreasing amounts of ethylene glycol (5.57%, 2.79% and 0% [v/v]). Follicles were then transferred to a dish of room temperature L-15 medium containing 10% SPS. The dish was put on a 37ºC warming plate for 10 minutes before transferring the follicles to maintenance medium. Approximately 4-6 rounds of cryopreservation were carried out for each seeding temperature and tissue type.
Fixation and immunofluorescence
Follicles to be imaged were fixed in 4% paraformaldehyde at 37°C for 1 hour followed by storage in wash buffer [38]. Follicles were stained for F-actin by incubation in rhodamine-phalloidin (1:50, Invitrogen) and 1 µg/ml Hoechst (Invitrogen) overnight at 4°C. Follicles were washed 3 x 10 minutes in wash buffer prior to mounting in buffer containing saline/glycerol (1:1) and sodium azide 25 mg/ml. In order to maintain the three-dimensionality of the follicles during imaging, glass shards and wax were placed beneath the glass coverslip to prevent compression.

Confocal microscopy and linescan analysis
Follicles were imaged on a Leica SP5 confocal microscope with a 40x oil objective. Full z-stacks images are comprised from 0.3 µm optical sections were completed for each follicle (n=10 for each treatment group). Using Leica analysis software, 4-pixel lines were drawn throughout the zona pellucida around the oocyte, in a 0.3 µm optical section, in order to determine the frequency (number of TZPs per 10 µm segment surrounding the oocyte). Three independent linescans were completed on five different sections per follicle (totaling 15 linescans per follicle).

Birefringence analysis
To quantify the maintenance of TZPs throughout the zona pellucida in non-cryopreserved and cryopreserved follicles, we used the Abrio Imaging System (CRi, Woburn, MA), which detects birefringent properties of cells. Birefringence is an intrinsic property of a sample that contains molecular order and can be detected by a polarized light microscope (pol-scope) and is measured by differences in polarized light as the retardance value [39]. The Abrio system was customized for our inverted Nikon Eclipse TE-300. Birefringence was analyzed using 20x and 40x Plan Flour DIC objectives. The Abrio software was capable of determining the orientation as well as the retardance value of the birefringence properties.

Microinjection and dye tracking
Twenty-five non-cryopreserved (Day 0 after isolation and Day 2 of culture) and 25 cryopreserved (Day 0 immediately after thaw and Day 2 of culture) C57BL/6J x CBA/Ca 2-3 layer secondary follicles were injected with 10% Lucifer Yellow CH, Lithium Chloride (Invitrogen, L12926) in EmbryoMax Injection Buffer (Chemicon; Millipore, Billerica, MA) as previously described [40, 41]. Injections into the oocyte were made using MicroJek EggJek injection needles (0.5 µm inner diameter) for 1 second at 120 hPa, and the dye was allowed to transfer to the surrounding granulosa cells via gap junctions for 10 minutes before being imaged on a Leica SP5 confocal microscope fitted with a resonance scanner. In order to assess the gap junction-TZP coupling, ten Lucifer Yellow fluorescent intensity measurements (0-255 units) were taken per follicle in the area of the granulosa cells. Two intensity measurements were taken every 5th section on 0.3 µm optical slices. The average intensity was then plotted against the average number of TZPs per 10 µm for each treatment and time point. All data was collected on and analyzed on a Leica TCS SP5 resonance scanning confocal microscope with Leica LAS AF software.

Statistical analysis
Transzonal projection counts and retardance measurements from control (non-cryopreserved) follicles and follicles pre- and post-cryopreservation and thaw were subjected to one-way
analysis of variance followed by Bonferroni’s multiple comparison post hoc-test to determine the significance of each treatment group using Prism 4 (GraphPad Software, Inc., La Jolla, CA). Calculated values are shown ±SEM with a significance level of P<0.05 being considered significant. Retardance measurements from follicles pre- and post-cryopreservation and thaw were plotted in box and whisker plots to represent the median value, minimum and maximum values and upper and lower quartiles.

**Results**

*Growth and survival of non-cryopreserved and cryopreserved mouse follicles*

As a control against which we could evaluate the successful cryopreservation and thawing of individual ovarian follicles, we first analyzed the growth characteristics of non-cryopreserved follicles grown in the 3D alginate hydrogel culture system. Two-layered secondary follicles were isolated and cultured for 10 days, and immunofluorescent images were taken every two days to identify healthy and unhealthy/dead follicles. After two days, 100% of cultured follicles were healthy (Figure 1A), with intact basement membranes and healthy, visible oocytes (not shown). By 10 days, 91.5±5.96% of follicles were healthy and had grown from ~150 µm to 319±11.1 µm in diameter (Figure 1B).

Since mouse follicles progress well to terminal maturation we used two seeding temperatures to more rigorously test the assays ability to detect the correlation between TZPs and follicle health. Follicles were cryopreserved at two seeding temperatures, -7ºC (permissive) and -9ºC (non-permissive), based on previously validated seeding temperatures for ethylene glycol in embryo and ovarian tissue cryopreservation [42-44]. After thawing, follicles were encapsulated in alginate and cultured for 10 days, with survival assessed every two days. It was found that the two-day survival rate for follicles seeded at the permissive temperature was approximately 91.6±0.60%, compared with the survival rate for follicles seeded at the non-permissive temperature (73.1±7.81%) (Figure 1A). At 10 days of culture, follicles seeded at -7ºC had a much higher survival rate (82.7±2.29%) than follicles seeded at -9ºC (67.5±6.67%) (Figure 1A). Accordingly, it was found that follicles seeded at -7ºC grew in diameter from an average of 153±7.38 to 330±11.7 µm, which was a statistically (P<0.01) larger increase than that seen in follicles seeded at the non-permissive temperature (from 133±3.52 to 255±22.2 µm; Figure 1B).

*Analysis of TZPs in cryopreserved mouse follicles*

Because follicle growth and survival is dependent on communication between the granulosa cells and the oocyte via TZPs, we determined whether TZPs survived the cryopreservation process or were capable of re-growth after cryopreservation and thawing. A selection of cultured follicles (both non-cryopreserved and cryopreserved) were fixed on Day 0, 2, 6 and 10 during the culture period and stained to examine the organization of TZPs in the area of the zona pellucida by confocal microscopy (Figure 2A).

Non-cryopreserved two-layer secondary follicles contained, on average, 8 TZPs per 10 µm of zona space surrounding the oocyte (Figure 2B). At two and six days of culture, TZPs in non-cryopreserved follicles increased to 11 per 10 µm, and by day 10, there were 12 TZPs per 10 µm (Figure 2B). Immediately thawed follicles showed few TZPs, with 4 and 2 TZPs per 10 µm for follicles seeded at -7 and -9ºC, respectively. At Day 2, TZPs increased to 9 (-7ºC) and 3 (-9ºC) TZPs per 10 µm. By 10 days of culture, TZPs in follicles seeded at the non-permissive
temperature increased to 8 TZPs, whereas TZPs in follicles seeded at -7°C increased to 11 per 10 µm section of zona (Figure 2B).

After 10 days of culture, follicles were stimulated for 16 hours to induce in vitro maturation, then the oocytes were examined for resumption of meiosis. It was found that 93% (n=67) of oocytes from non-cryopreserved follicles, 57.3% (n=35) of oocytes in the -7°C group and 30.7% (n=21) of oocytes from the -9°C group matured to the MII stage. MII staged oocytes were subjected to in vitro fertilization and were examined for pronuclear formation after 6 hours. It was found that 79% (n=53) of oocytes from the non-cryopreserved group, 57% (n=20) from the -7°C group and 52% (n=11) from the -9°C group were able to be fertilized and form pronuclei.

**Functional transzonal projection studies**

To assess the functionality of the TZPs in non-cryopreserved and cryopreserved mouse follicles (two hours after thawing), Lucifer Yellow was injected into the oocytes of non-cryopreserved and cryopreserved follicles and the ability of the dye to transfer from the oocyte to the surrounding granulosa cells through intact gap junctions was examined. After 10 minutes, the follicles were imaged for dye transfer by confocal microscopy (Figure 3). Injected non-cryopreserved follicles contained brightly stained oocytes surrounded by several stained granulosa cells indicative of intact gap junctions (Figure 3A, a). After 2 days of culture, isolated, non-cryopreserved follicles were injected and shown to transfer dye to surrounding somatic cells (Figure 3B, b). In cryopreserved follicles, immediately after thawing, no dye transferred from the oocyte to the surrounding granulosa cells (Figure 3C, c). However, after two days in culture, follicles cryopreserved at -7°C were injected and were able to transfer Lucifer Yellow dye to surrounding granulosa cells indicating the reformation of TZPs and gap junctions (Figure 3D, d). Very little dye transferred in follicles cryopreserved at a seeding temperature of -9°C after 2 days of culture (3E, e).

Though it was determined that TZPs were re-established and functional by the passage of Lucifer Yellow dye from the oocyte to the granulosa cells in -7°C follicles it was necessary to establish the coupling strength of the gap junctions at the ends of TZPs in these follicles (Figure 4). The amount of dye transferred to the granulosa cells was quantified by the average fluorescent intensity per 0.3 µm section and was plotted against the average number of TZPs per 0.3 µm section. It was found that non-cryopreserved follicles were moderately coupled on the day of isolation having an average of 9 TZPs and an average fluorescent intensity of 60 units. However, after 2 days in culture, the TZPs and gap junctions were highly coupled increasing to 11 TZPs and 110 fluorescent units. Analysis of thawed follicles showed very little initial coupling of gap junctions and TZPs. Interestingly, after 2 days in culture follicles cryopreserved with a seeding temperature of -7°C increased in gap junction-TZP coupling where follicles cryopreserved with a seeding temp of -9°C did not. Overall, the number of TZPs directly correlated with the average fluorescent intensity of dye.

**Analysis of TZPs using birefringence analysis**

In order to avoid terminal assessment of follicle health, we wanted to test whether non-invasive polarized light is capable of detecting TZPs vitally in growing follicles, as a biomarker of follicle health. Therefore, we investigated an alternative technique of birefringence analysis, which is used to detect organized microtubules and F-actin in living cells [39]. We first investigated
whether a pol-scope was able to detect the organized proteins of the zona pellucida and TZPs by comparing the birefringence of the zona pellucida region of untreated mouse follicles (Figure 5A) and of follicles treated with latrunculin B, an actin depolymerizer (Figure 5B). In the untreated follicles, birefringence is seen around the oocyte in Figure 5A (inset-birefringence alone) and is absent from the images of follicles treated with actin depolymerizers for 5 hours (Figure 5B; inset-birefringence alone). Birefringence measurements in the area of the zona were recorded, and revealed that treating follicles with latrunculin B significantly decreased the retardance to 1.3 nm compared with 2.25 nm in untreated follicles (Figure 5E). Untreated and latrunculin B-treated follicles were fixed and prepared for confocal microscopy to image TZPs. Control follicles showed intact TZPs throughout the zona pellucida (Figure 5C), which had approximately 8.9±0.25 TZPs per 10 µm. As expected, follicles treated for 5 hours with latrunculin B displayed depolymerized actin bundles in the area of the zona (Figure 5D), containing 3.6±0.37 TZPs per 10 µm. From this assessment, we can conclude that in mouse follicles, birefringence of the zona pellucida occurs at levels <1.0 nm and signals >1.0 nm represent birefringence of TZPs.

We then analyzed the birefringence properties of non-cryopreserved and cryopreserved follicles during culture in alginate. All two-layer secondary follicles recorded similar retardance levels prior to freezing, ranging from 1.75 to 2.2 nm (Figure 6). Non-cryopreserved follicles continued to increase in birefringence throughout the six days of culture to 2.75 nm, consistent with the increase in TZPs during this culture period (Figure 2B). After thawing, day zero cryopreserved follicles showed a decrease in retardance to 1.0 nm; however, by day two of culture, follicles cryopreserved at the permissive seeding temperature showed an increase to 2.0 nm, consistent with the re-growth in TZPs seen in Figure 2B, and by day six, these follicles increased retardance to 2.5 nm. However, follicles cryopreserved at the non-permissive seeding temperature failed to significantly increase levels of retardance throughout the six-day culture period, ending at 1.25 nm (Figure 6).

**Cryopreservation of non-human primate follicles**
To determine whether cryopreservation of individual follicles can be applied to human fertility preservation, and to test whether the birefringence assay is adaptable to assessing follicle health in other species, we isolated individual secondary follicles from rhesus macaques and analyzed their follicle growth characteristics in alginate culture as well as the birefringent properties of their TZPs.

Isolated follicles were less than 200 µm in diameter and increased to 350 µm during six days of culture (Figure 7A). Interestingly, the birefringence properties of non-cryopreserved rhesus follicles were consistent throughout the growth period, ranging from 3 to 4 nm (Figure 7B). Cryopreserved follicles (using the permissive seeding temperature) were approximately 175 µm in diameter after thawing and grew to over 250 µm during six days of culture (Figure 7C). The birefringent properties of cryopreserved rhesus follicles could be detected using a pol-scope, with pre-cryopreservation retardance values of approximately 2.75 nm. Following thaw (post-cryopreservation), values decreased on average to 1.25 nm. By two days of culture, however, the retardance values increased to approximately 3.0 nm, which was maintained throughout the six-day culture period (Figure 7D), consistent with that seen in Figure 7B. On day six, both non-
cryopreserved and cryopreserved follicles were fixed and assessed for TZPs; they contained 17.1 ± 0.51 (Figure 7E) and 14.3 ± 0.83 TZPs per 10 μm, respectively.

Cryopreservation of human follicles

It has been established that TZPs are reflective of birefringence in mice (Figure 5) and rhesus follicles show numerous TZPs (Figure 7E), which associate with high levels of birefringence (Figure 7B). Therefore, we lastly examined the health of non-cryopreserved and cryopreserved follicles from four human oncofertility patients to determine if birefringence from human follicles, as shown by the bright birefringent ring in the area of the zona (Figure 8A), positively correlates with follicle growth and health. Non-cryopreserved follicles were grown for six days in the alginate culture system. Diameter and birefringence were plotted for each individual follicle (Figure 8C). Follicle sizes ranged from 80-425 μm in diameter with retardance ranging from 1.5 to 5.25 nm. Non-cryopreserved follicles were found to have a positive correlation between increased follicle diameter and high levels of birefringence. Though cryopreserved human follicles displayed little birefringence immediately following thaw (Figure 8B), follicles were capable of reestablishing physical connections. Data from cryopreserved follicles (-7ºC seeding temp) were also plotted according to follicle diameter and retardance value (Figure 8D). Follicles ranged from 80-175 μm in diameter having retardance values from 0.75-2.5 nm. Cryopreserved human follicles were also found to have a positive correlation between follicle growth and an increase in birefringence (Figure 8D). On day six of culture, fixed-follicle analysis revealed that non-cryopreserved and cryopreserved follicles contained 11.3 ± 0.34 and 7.0 ± 0.58 TZPs per 10 μm, respectively.

Discussion

Understanding the impact of cryopreservation of secondary ovarian follicles not only allows researchers an opportunity to study the importance of granulosa cell-oocyte communication for follicle survival, it also holds potential for expanding fertility preservation options for women facing fertility-threatening diseases or treatments, and improving methods of preserving tissue of endangered species. Various groups have studied cryopreservation of ovarian tissue as well as the preservation of oocytes and embryos; however, we believe this is the first study to examine the importance of preserving the connectivity between the granulosa cells and the oocyte after cryopreservation. Here we report that 1) individual secondary follicles (mouse, rhesus monkey and human) may be cryopreserved and grown in a 3D matrix; 2) successful follicle growth following cryopreservation correlates with the ability of follicles to reestablish and remodel TZPs; and 3) vital detection of TZPs via birefringence is a reflection of follicle health and can be detected following cryopreservation.

Follicle growth and survival post-cryopreservation

Secondary follicles in situ fail to survive the cryopreservation process due to ice formation in the large fluid volumes of the oocyte and follicle, as well as poor penetration of the cryoprotectant [45]. We hypothesized that isolation from ovarian cortical tissue and cryopreservation of individual preantral secondary follicles would increase the follicle survival rate after thawing. Our group has previously demonstrated the ability to culture mouse [21], rhesus macaque [22] and human [23] secondary follicles in 3D alginate hydrogels, which is an essential step for resuming follicle growth after the cryopreservation process. Using this alginate hydrogel as a 3D matrix for follicle growth, we were able to assess the follicle health and growth following
cryopreservation. Using a permissive (-7°C) seeding temperature for ethylene glycol, follicles survived the cryopreservation process and grew in culture for 10 days to a size comparable to non-cryopreserved cultured follicles. We were also able to induce oocyte maturation in situ and fertilize MII oocytes from individual cryopreserved follicles, indicating that the zona pellucida did not lose its ability to interact and bind sperm. Rhesus monkey and human follicles were also individually cryopreserved and their growth characteristics were compared to non-cryopreserved follicles. Although it was found that both rhesus monkey and human cryopreserved follicles grew slower following cryopreservation, they continuously increased follicle diameter during the entire culture period.

Transzonal projection remodeling in cryopreserved follicles
In addition to being critical for bidirectional communication, it has been reported that TZPs play a role in maintaining the organization of the oocyte cortex in mice [46]. TZPs, which are primarily made of F-actin and form during the course of follicle development, are extremely sensitive to fluctuations in temperature [47-49] and are therefore as vulnerable as oocytes with regard to the destruction of the microtubule and actin network during the cryopreservation process [50]. Therefore, to best protect secondary follicles from the adverse cryo-effects on microtubules and microfilaments, follicles were individually isolated for better permeation of cryoprotectant.

Our confocal analysis of TZPs after cryopreservation showed that a large percentage of the physical connections in mouse follicles were destroyed after the thawing process but that they were capable of reconnecting in the 3D culture system; TZPs increased in number as the follicle increased in diameter during the first six days of culture. Follicles that were cryopreserved at the non-permissive seeding temperature (-9°C) were unable to reestablish and maintain connectivity following cryopreservation, ultimately resulting in follicle death (seen in Figure 1A). This highlights the importance of seeding temperature in the slow cryopreservation process. TZPs were also shown to be functional by the injection and transport of Lucifer Yellow from the oocyte into the surrounding granulosa cells over two days of culture. Immediately after thawing, follicles were unable to transfer dye into surrounding granulosa cells, indicating that either TZPs were not present or that gap junctions were not functional. However, after two days of culture, cryopreserved follicles (-7°C) were able to re-establish TZPs and gap junctions and transport dye from the oocyte into the surrounding somatic cells. This is the first report to establish that TZPs are able to re-grow and function in follicles that have been cryopreserved.

Birefringence analysis
Oocyte and embryo quality are assessed, both in the laboratory and the clinic, by several visual characteristics, such as the placement of the spindle and polar body and cleavage rate of blastomeres [51, 52]. However, the only way to vitally assess follicle health is by monitoring follicle growth and hormone secretion over several days [21-23]. We tested the effectiveness of an alternative vital assay of follicle health that uses a pol-scope to detect birefringence in the area of TZPs (zona pellucida). Microtubules and actin filaments, which are predominant components of TZPs, are organized structures that are easily detected by the pol-scope, similar to the microscopes used to detect the MII spindle in fertility clinics and transgenic labs [53-55].
We first tested the effectiveness of the polarized-light microscope to detect TZPs by depolymerizing actin filaments in follicles using latrunculin B. Treated follicles had a reduced number of TZPs per 10 µm and a concomitant reduction in birefringence in the area of the TZPs, a phenomenon that was not seen in untreated control follicles. It was established that the zona pellucida has an intrinsic birefringence level of 1.0 nm; additional birefringence above 1.0 nm is contributed by TZPs. Using the pol-scope on both non-cryopreserved and cryopreserved growing mouse follicles in culture, the number of TZPs increased and decreased (Figure 2B) in correlation with increases and decreases in birefringence (Figure 6). The assessment of rhesus monkey and human follicles further validated the effectiveness of this vital assay on measuring the health of follicles (Figures 7 and 8). It was found that both non-cryopreserved and cryopreserved rhesus monkey follicles grew over a six-day culture period and maintained high levels of birefringence, even after cryopreservation, which was consistent with high TZP counts seen on the last day of culture. Non-cryopreserved and cryopreserved human follicles were assessed for health by examining follicle diameter and retardance values during six-days of culture. Both non-cryopreserved and cryopreserved follicles indicated that follicle size directly correlates with an increase in retardance, which is reflective of follicle health. Though cryopreserved follicles showed less of an increase in follicle size and birefringence over the course of the six-day culture period, which may be a result cryopreservation, we are still able to vitally assess follicle health using this method. This bioassay provides the badly needed technique that can immediately, systematically and non-invasively monitor early follicle health after a cryopreservation insult. As new cryo-technologies emerge, this method provides a signature endpoint that can be exploited routinely.

Our studies reveal that mouse, monkey and human follicles are capable of surviving current cryopreservation techniques and that the follicle health can be evaluated by a non-invasive imaging technique that detects intact TZPs. This technique may permit selection of healthy cyropreserved follicles that can then be encapsulated in a 3D alginate hydrogel matrix and grown in culture to produce healthy oocytes. By enriching the cultured follicle population in this way, we would expect to achieve higher rates of follicle survival and growth as well as higher rates of oocyte maturation and fertilization. The next challenge will be to determine how long non-human primate and human follicles can be cultured after cryopreservation, and to incorporate our vital assay into the 30-day primate and human follicle culture protocol [22, 23] to assess follicle and oocyte health over time.

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**References**


Figure Legends
Figure 1. (A) Survival in non-cryopreserved (Non-cryo) and cryopreserved (with seeding temperatures of -7 ºC and -9 ºC) follicles over a 10-day culture period. Error bars represent ±SEM. *P<0.05. (B) Growth of non-cryopreserved and cryopreserved follicles between day two and day 10 of culture. Error bars represent ±SEM. *P<0.01.
Figure 2. (A) TZP frequency is recorded for every 10 µm around the follicle perimeter in the area of the zona pellucida (*). F-actin (purple), DNA (blue-green). Magnification bar represents 20 µm. Inset shows an enlarged area of the zona depicting individual TZPs from one 0.3 µm section of the full Z-stack (B). Arrows depict individual TZPs. TZP frequency in non-cryopreserved and cryopreserved cultured mouse follicles over a 10-day culture period. Error bars represent ±SEM. *P<0.001, **P<0.01.

Figure 3. Microinjection of Lucifer Yellow tracking dye to detect functional TZPs and gap junctions in day 0 (A, a) and day 2 (B, b) non-cryopreserved follicles and day 0 (C, c) and day 2 (D, d) cryopreserved follicles seeded at -7ºC and day 2 (E, e) cryopreserved follicles seeded at -9ºC. Brightfield images are shown in lower case panels. Magnification bar represents 25 µm (A, a, C, c), 15 µm (B, b) and 20 µm (D, d). (F) Ratio of TZP density over the average fluorescent intensity after dye transfer on day 0 and day 2 of culture of cryopreserved and non-cryopreserved follicles.

Figure 4. Coupling strength of TZPs to gap junctions in non-cryopreserved and cryopreserved follicles. The average TZP number per 10 µm and the average fluorescent intensity of Lucifer Yellow in the granulosa cells shows that gap junctions are highly coupled in non-cryopreserved follicle. Immediately after thawing, gap junctions and TZPs are not coupled in -7 and -9ºC. However after 2 days in culture gap junction-TZP coupling have increased in -7ºC yet -9ºC have not. Error bars represent ±SEM. (a) Represents P<0.01 between -7 and -9ºC on the same day. (b) Represents P<0.01 between -7 and P<0.001 between -9ºC on same day. (c) Represents P<0.01 between -9ºC on same day.

Figure 5. Birefringence analysis of TZPs in follicles treated with the actin depolymerizing drug latrunculin B. Birefringent images of untreated follicles (A) and follicles after five hours of latrunculin B (Lat B) treatment (B). Insets represent birefringence of the cell marked with (*). Confocal image of a five-hour control follicle stained for F-actin (green) to show transzonal projections (C, inset). Confocal image of a follicle treated for five hours with latrunculin B and stained for F-actin (green) showing no transzonal projections (D, inset). (E) Non-treated follicles have a much higher retardance than follicles treated for five hours with 250 µM latrunculin B. A retardance score below 1.0 nm represents birefringence from the zona and a retardance score above 1.0 nm represents birefringence from TZPs. Error bars represent ±SEM. Magnification bar represents 25 µm.

Figure 6. Analysis of the birefringent properties of non-cryopreserved and cryopreserved mouse follicles cultured for six days. Fresh follicles show an increasing retardance over six days of culture. Cryopreserved follicles showed a decrease in retardance after cryopreservation at the permissive temperature but an increase back to non-cryopreserved levels after 2 days in culture. Error bars represent ±SEM. *P<0.01, **P<0.001 between treatment groups on the same day.

Figure 7. Growth and birefringent properties of non-cryopreserved and cryopreserved rhesus monkey follicles. (A) Increased growth of non-cryopreserved follicles during six days in culture. (B) The birefringent properties of fresh follicles did not change during six days in culture and maintained retardance values between 3 and 4 nm. (C) Growth of cryopreserved follicles after six days in culture. (D) Birefringence of cryopreserved follicles showed a decreased retardance.
after cryopreservation and an increased retardance to non-cryopreserved levels after two days in culture. (E) Staining of TZPs (green) in a non-cryopreserved rhesus follicle, DNA (red). Box plot graphs represent the median value and minimum and maximum values as well as the upper and lower quartiles. Mean ± SEM marked with different superscript letters are significantly different (P<0.05) between culture days. Magnification bar represents 20 µm (E).

Figure 8. Growth and birefringent properties of non-cryopreserved and cryopreserved human follicles. (A) Birefringent image of a human follicle grown for 6 days in alginate hydrogel; high levels of retardance are seen in the area of the zona. (B) Birefringent image of a human follicle: note the lower levels of retardance are seen in the area of the zona. Colors represent directionality of the actin fibers. Magnification bar represents 30 µm (A) and 50 µm (B). (C) Analysis of TZP retardance and follicle diameter in non-cryopreserved follicles and (D) cryopreserved follicles. Regression lines show a positive trend for both non-cryopreserved (y = 0.0095x + 1.183, R² = 0.3913) and cryopreserved follicles (y = 0.0085x + 0.5512, R² = 0.15243).
Figure 1

A

% Survival

Day 2   Day 10

B

Follicle Diameter (μm)

Day 2   Day 10
Figure 2